(19) World Intellectual Property Organization

International Bureau



(43) International Publication Date 26 August 2004 (26.08.2004)

PCT

(10) International Publication Number WO 2004/071421 A2

(51) International Patent Classification7:

A61K

Street 10, 69018 Tel Aviv (IL), LEVY, Ruth [IL/IL]; Sasha Argov Street 25, Apt. #2, 69620 Tel-Aviv (IL).

(21) International Application Number:

PCT/US2004/003281

(22) International Filing Date: 5 February 2004 (05.02.2004)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

60/445,328

5 February 2003 (05.02.2003) US

- (71) Applicant (for all designated States except BB, US): TEVA PHARMACEUTICAL INDUSTRIES LTD. [IL/IL]; 5 Basel Street, P.O. Box 3190, Petah Tikva 49131 (IL).
- (71) Applicant (for BB only): TEVA PHARMACEUTICAL USA, INC. [US/US]; 1090 Horsham Road, North Wales, PA 19454 (US).
- (72) Inventors; and
- (75) Inventors/Applicants (for US only): LICHT, Daniella [II]/II.]; Keren Ha Yesod 1, 54051 Ramat Han Givat Shmeul (IL). LOVINGER, Ioana [IL/IL]; Rh. Tlamin 15/3, Ganci-Aviv., 71338 Lod (IL). ABD-ELHAI, Suher [IL/IL]; 44915 Tira (IL). DAGAN-LION, Mazzi [IL/IL]; Nahal netafim 22, 71700 Modiin (IL), GILBERT, Adrian [IL/IL]; 60B Apt., Hankin Street 5, 43465 Ra'anana (IL). LEIBOVITCH, Noa [II/IL]; Nachshon St. 49, 47301 Ramat-Ha Sharon (IL). COHEN, Sasson [IL/IL]; Meyzan

(74) Agent: WHITE, John, P.; Cooper & Dunham LLP, 1185 Avenue of the Americas, New York, NY 10036 (US).

- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GD, GM, HR, HU, HJ, H., IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.
- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GII, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: SUSTAINED RELEASE FORMULATION OF N-(2-PROPYLPENTANOYL) GLYCINAMIDE AND RELATED COMPOUNDS

(57) Abstract: The subject provides a sustained release tablet comprising the following components: a) a uniform admixture of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure: (I) wherein R₁, R₂, and R₃ are independently the same or different and are hydrogen, a C₁-C₆ alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and a binder, and b) a hydroxypropylmethyl cellulose, a process for manufacturing the tablet and a manufacturing the tablet and a manufacturing the companion of effecting pain prophylaxis



SUSTAINED RELEASE FORMULATION OF N-(2-PROPYLPENTANOYL)GLYCINAMIDE AND RELATED COMPOUNDS

This application claims the benefit of U.S. Provisional Application No. 60/445,328, the entire contents of which are hereby incorporated by reference.

10 Throughout this application, various publications are referenced by full citations. The disclosures of these publications in their entireties are hereby incorporated by reference into this application in order to more fully describe the state of the art as known to those skilled therein as of the date of the invention described and claimed herein.

Background of the Invention

5

Pain is considered to play a basic physiological role in the detection and localization of tissue damage or potentially damaging physiological processes. Pain has been broadly classified as somatogenic, where a physiological explanation can be found, or psychogenic, where the physiological explanation is not known (The Merck Manual of Diagnosis and Therapy, 16th Ed., pp. 1407-1426; PCT International Publication No. WO 02/13766 A2). An example of somatogenic pain is neuropathic pain.

Neuropathic pain is a category of pain which includes several forms of non-nociceptive chronic pain, which result from dysfunction of nervous rather than somatic tissue. The majority of non-nociceptive chronic pains, in terms of either syndromes or cases, follow at various times after damage to either central or peripheral nervous tissue. Diagnosis of most of these syndromes and cases reveals a dependence on abnormal spatial and temporal summation of natural somatic stimulation

in the spinal cord and independence from somatic disease and peripheral sympathetic nervous system activity. The scientific pain research community defines this kind of pain as centrally mediated neuropathic pain and recognizes mechanistic, diagnostic, and therapeutic commonalities among pains of this class and differences between these and other syndromes.

Neuropathic pain can be defined as pain deriving from damage to or inflammation of central or peripheral nervous system tissue.

10 Examples of pain syndromes of this class include post herpetic neuralgia, neuritis, temporomandibular disorder, myofascial pain, back pain, pain induced by inflammatory conditions. Neuropathic pain may occur in all body regions. For example, neuropathic pain may originate from the dental region.

15

Burn injury also often leads to neuropathic hyperalgesia in the affected body area. Neuralgia is characterized, in its acute phase, by intraneural inflammation, which can cause damage to primary afferent axons, thus inducing neuropathic pain.

Neuropathic pain may also be induced by diabetic conditions (diabetic neuropathy). Neuropathy of primary afferent axons in long nerves is found in diabetic patients. Nociceptor

sensitization may ensue (U.S. Patent No. 6,054,461).

25 Pain can be both chronic and acute, and can also be evoked by noxious stimuli, referred to as hyperalgesia, or by non-noxious stimuli, referred to as allodynia (Attal, N. "Mechanism of action and rationale for use of antiepileptic drugs" (1999) in International Congress and Symposium Series 241 The Royal 30 Society of Medicine Press, Limited Ed. JM Pellock). Allodynia and hyperalgesia can have mechanical causes (dynamic or static), or a thermal cause. Examples of neuropathic pain peripheral neuropathies include all the painful postherpetic specifically diabetic peripheral neuropathy, 35 neuralgia, and trigeminal neuralgia. Trigeminal neuralgia, for

example, is the most common neuralgic syndrome in the elderly.

Other types of somatogenic pain that may have neuropathic components include cancer pain, postoperative pain, lower back pain, complex regional pain syndrome, phantom pain, HIV pain, arthritis (osteo-arthritis and rheumatoid arthritis) pain and migraines.

Pain may also be a symptom of headache disorders. Migraines constitute one of the four major categories of primary headaches (International Headache Society, 1988; Silberstein, S.D. et al. <u>Headache in Clinical Practice</u>, (1998) Pub. Isis Medical Media, Oxford). The other three types of primary headaches are tension -type, cluster and a miscellaneous-type (Id.). One current view is that there is a continuous spectrum of headache severity ranging from mild tension headaches to severe migraines. Others consider tension headaches and migraines to be distinct entities.

Neuropathic pain conditions are characterized by hyperesthesia

(enhanced sensitivity to a natural stimulus), hyperalgesia
(abnormal sensitivity to pain), allodynia (widespread
tenderness, characterized by hypersensitivity to tactile
stimuli), and/or spontaneous burning pain.

- 25 The initial drug of choice for treating trigeminal neuralgia is carbamazepine. For other types of pain, such as postherpetic neuralgia and painful diabetic neuropathy, amitriptyline is most commonly used.
- 30 Drugs used in the treatment of headache disorders such as migraines originate from a broad range of different drug categories. These include: 5-hydroxytryptamine agonists (5-HT₁ agonists), dihydroergotamine, ergotamine, anti-emetics, anxiolytics, non-steroidal anti-inflammatory drugs, steroids, 35 major tranquilizers, narcotics, beta-blockers, calcium channel

Ø

blockers, anti-depressants, and anti-epileptic drugs. However, not all of the drugs in these categories are truly effective. There is still a need for more efficacious drugs, as well as a need for antimigraine treatments with fewer side effects.

Epilepsy is an ancient disease, which affects about 1% of the global population. Despite the progress made in antiepileptic drug therapy, there are still many patients who continue to suffer from uncontrolled seizures and medication toxicity. At present, only the following 4 major antiepileptic drugs are in use: phenobarbital, phenytoin, carbamazepine and valproic acid. About 25% of the patient population is not seizure-free while treated with these medications (both mono and polytherapy) even when diagnosis and therapy is optimal ("Sustained Release Formulations of Antiepileptics" Clin. Pharmacokinet. (1992) 22(1): 11-24).

Pharmacological activity in general and antiepileptic activity in particular, correlate better with the concentration of a drug in the blood (or in some other biophase) than with the administered dose. This phenomenon is due, in part, to variability in drug absorption and disposition between and within individuals, particularly when the drug is given orally.

Optimizing drug therapy aims at achieving and maintaining therapeutic and safe drug concentration in the blood. In order to achieve this goal, it would be advantageous, and probably more convenient, that the patient receive a once- or twice-daily dosage regimen (Ballard 1978; Silber et al. 1987, Welling 1983).

N-(2-Propylpentanoyl)glycinamide is an anti-epilepsy and antipain drug which has the structure:

and can be prepared as disclosed by Bialer et al. in U.S. Patent 5,585,358. U.S. Patent 5,585,358 also describes a series of derivatives of valproic acid amides and 2-valproenic acid amides for the treatment of epilepsy and other neurological disorders.

Bialer et al. refer to the above compound as N-(2-n-Propylpentanoyl)glycinamide. However, in the present application, the compound is referred to as N-(2-Propylpentanoyl)glycinamide.

Published U.S. Patent Application No. US-2002-0052418-Al discloses the use of N-(2-Propylpentanoyl)glycinamide and other derivatives of valproic acid amides and 2-valproenic acid amides for the treatment or prevention of pain and/or headache disorders.

U.S. Patent 5,009,897, issued April 23, 1991 discloses granules, suitable for pressing into tablets, the granules comprising a core of divalproex sodium and a coating of a mixture of a polymer and microcrystalline cellulose.

U.S. Patent 4,913,906, issued April 3, 1990, discloses controlled release dosage forms of valproic acid, its amide, or one of its salts or esters in combination with a natural or synthetic polymer, pressed into a tablet under high pressure.

U.S. Patent 4,913,906 does not, however, disclose the use of hydroxypropylmethyl cellulose, or the use of two or more materials to achieve controlled release.

U.S. Patent 6,419,953, issued July 16, 2002, discloses controlled release formulations of valproic acid, its salt, divalproex sodium, or valpromide, comprising granules of the 5 active ingredient, each granule containing the active compound, and lactose, \mathtt{mixed} hydroxypropylmethyl cellulose In U.S. Patent 6,419,953, additional excipients. hydroxypropylmethyl cellulose, if used, is part of each granule. U.S. Patent 6,419,953 does not disclose compressing active ingredient with hydroxypropylmethyl of 10 granules cellulose.

The subject invention provides a sustained release formulation of N-(2-Propylpentanoyl) glycinamide.

Summary of Invention

5

10

15

20

The subject provides a sustained release solid dosage form comprising the following components:

a) a uniform admixture of:

(i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

a compound having the
$$N$$
 of N of

or $(CH_2)n$ NR_2R_3

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

(ii) a binder, and

b) a hydroxypropylmethyl cellulose.

25 The subject invention also provides a sustained release tablet comprising the following components:

- a) a uniform admixture of:
 - (i) N-(2-Propylpentanoyl)glycinamide; and
 - (ii) a binder;
- b) a hydroxypropylmethyl cellulose; and
- c) a different hydroxypropylmethyl cellulose.

The subject invention also provides a hard compressed tablet comprising a uniform admixture of the following components:

- a) N-(2-Propylpentanoyl)glycinamide;
- b) a hydroxypropylmethyl cellulose; and
- c) a different hydroxypropylmethyl cellulose.

The subject invention also provides a composition in granulate form comprising a uniform admixture of:

(i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N \in \mathbb{N}} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

25

20

5

10

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

(ii) a hydroxypropyl cellulose.

Detailed Description of the Figures

Figure 1 shows mean plasma N-(2-propylpentanoy1) glycinamide concentrations following the administration of 2 x 500mg N-(2-propylpentanoy1) glycinamide tablets (Formulation A), N-(2-propylpentanoy1) glycinamide tablets (Formulation B) and 2 x 500mg N-(2-propylpentanoy1) glycinamide tablets (Formulation C) to eighteen healthy male Caucasian volunteers.

- 10 -•- Formulation A
 - -o- Formulation B
 - -*- Formulation C
- Figure 2 shows mean plasma concentrations of N-(2-15 propylpentanoyl) glycine following the administration 2 500mg N-(2-propylpentanoyl) glycinamide 500 tablets (Formulation A), 2 x N-(2propylpentanoyl) glycinamide tablets (Formulation B) and 2 x 500mg N-(2-propylpentanoy1) glycinamide . 20 tablets (Formulation C) to eighteen healthy male Caucasian volunteers.
 - --- Formulation A
 - -0- Formulation B
 - -*- Formulation C

Detailed description

The subject invention provides a sustained release solid dosage form comprising the following components:

- a) a uniform admixture of:
 - (i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

or.

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

15

10

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

(ii) a binder; and

25

20

b) a hydroxypropylmethyl cellulose.

In one embodiment, the solid dosage form is a tablet.

In another embodiment, the uniform admixture of component a) further comprises a filler.

In one embodiment, the filler comprises a microcrystalline cellulose.

In another embodiment, the hydroxypropylmethyl cellulose comprises 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxyproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve, has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C, and has a pH in the range 5.5-8.0.

In another embodiment, at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

20 In another embodiment, the solid dosage form further comprises as additional components a filler, a lubricant and a flow agent.

In another embodiment, the binder of component a)(ii) comprises

25 hydroxypropyl cellulose.

In another embodiment, the solid dosage form further comprises a different hydroxypropylmethyl cellulose as a component.

30 In another embodiment, the solid dosage form further comprises as additional components a filler, a lubricant and a flow agent.

In another embodiment, the solid dosage form further comprises a different hydroxypropylmethyl cellulose.

In embodiment, the different hydroxypropylmethyl cellulose comprises 19-24% by weight methoxyl substituent, 7-9% weight hydroxypropoxyl substituent, has an 5 viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C, has a pH in the range 5.5-8.0 and has a particle size distribution such that at least 99% of 10 hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

In another embodiment, at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

15

20

25

In another embodiment,

the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose, methylcellulose, carboxymethylcellulose, calcium carbonate, calcium sulfate kaolin, sodium chloride, powdered cellulose, sucrose, mannitol, starch, corn starch, various natural gums or a combination of two or more of the foregoing;

the lubricant comprises magnesium stearate, sodium stearyl fumarate, hydrogenated castor oil, hydrogenated soybean oil, polyethylene glycol or a combination of two or more of the foregoing; and

the flow agent comprises a colloidal fumed silica, or colloidal silicon dioxide.

30 In another embodiment,

the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant comprises magnesium stearate, sodium stearyl fumarate or a combination thereof; and

the flow agent comprises a colloidal fumed silica.

In another embodiment, the active ingredient is a compound having the structure:

or

$$\bigcap_{H} (CH_2)n \bigcap_{NR_2R_3}$$

10

5

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3.

15

25

In a further embodiment, the active ingredient is N-(2-Propylpentanoyl)glycinamide.

In another embodiment, the above solid dosage form also comprises the following components:

- a) a uniform admixture of:
 - (i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide, N-(2-Propylpentanoyl)glycinamide,

N-(2-propylpentanoyl)glycine-N'-methylamide,

N-(2-propylpentanoyl)glycine-N'-butylamide,

N-(2-propylpentanoyl)leucinamide,

N-(2-propylpentanoyl)alanine-N'-benzylamide,

N-(2-propylpentanoyl)alapinamide,

N-(2-propylpentanoyl)-2-phenylglycinamide,

N-(2-propylpentanoyl) threoninamide,

N-(2-propylpentanoyl)glycine-N', N'-dimethylamide,

N-(2-propylpent-2-enoyl)glycinamide,

N-(2-propylpent-2-enoyl)alaninamide, and

N-(2-propylpent-2-enoyl)glycine-N'-methylamide; and

- (ii) a binder, and
- b) a hydroxypropylmethyl cellulose.

15 The subject invention also provides a sustained release solid dosage form comprising the following components:

- a) a uniform admixture of:
 - (i) N-(2-Propylpentanoyl)glycinamide; and
 - (ii) a binder;
- 20 b) a hydroxypropylmethyl cellulose; and
 - c) a different hydroxypropylmethyl cellulose.

In one embodiment, the solid dosage form is a tablet.

- In one embodiment, the solid dosage form comprises a filler, a lubricant and a flow agent as additional components and wherein the uniform admixture of component a) further comprises a filler.
- 30 In another embodiment,

5

10

the binder of component a) (ii) comprises hydroxypropyl cellulose;

the filler of component a) comprises a microcrystalline cellulose;

the hydroxypropylmethyl cellulose of component b) has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;

the hydroxypropylmethyl cellulose of component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;

the filler component comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant component comprises magnesium stearate, sodium stearyl fumarate or a combination thereof; and

the flow agent component comprises a colloidal fumed silica.

In another embodiment, the solid dosage form comprises the following components:

a) a uniform admixture of:

5

10

15

25

- (i) from 50 mg/solid dosage form to 1000 mg/solid dosage form of N-(2-propylpentanoyl)glycinamide,
- (ii) from 1 mg/solid dosage form to 100 mg/solid
 dosage form hydroxypropyl cellulose; and
 (iii) from 1 mg/solid dosage form to 200 mg/solid
 dosage form microcrystalline cellulose;
- b) from 10 mg/solid dosage form to 300 mg/solid dosage form of hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 78-128 by hydroxylproproxyl substituent and has a particle size distribution at such that least 998 the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
- 35 c) from 10 mg/solid dosage form to 300 mg/solid dosage form of a different hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12%

hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;

- d) from 1 mg/solid dosage form to 300 mg/solid dosage form microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
 - e) from 0.1 mg/solid dosage form to 20 mg/solid dosage form of magnesium stearate, sodium stearyl fumarate or a combination thereof; and
 - f) from 0.1 mg/solid dosage form to 15 mg/solid dosage form a colloidal fumed silica.
- In another embodiment, the solid dosage form comprises the following components:
 - a) a uniform admixture of:

10

25

30

- (i) from 500 mg/solid dosage form to 850 mg/solid dosage form of N-(2-propylpentanoy1)glycinamide,
- 20 (ii) from 25 mg/solid dosage form to 75 mg/solid dosage form hydroxypropyl cellulose; and
 - (iii)from 50 mg/solid dosage form to 150 mg/solid
 dosage form microcrystalline cellulose;
 - b) from 100 mg/solid dosage form to 300 mg/solid dosage form of hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
 - c) from 20 mg/solid dosage form to 150 mg/solid dosage form of a different hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the

hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;

- d) from 20 mg/solid dosage form to 100 mg/solid dosage form microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
- e) from 2 mg/solid dosage form to 20 mg/solid dosage form of magnesium stearate, sodium stearyl fumarate or a combination thereof; and
- f) from .5 mg/solid dosage form to 5 mg/solid dosage form a colloidal fumed silica, per 1 gram solid dosage form.

In one embodiment, at least 90% of the hydroxypropylmethyl cellulose of component b), of component c), or of both component b) and c) passes through a No. 100 US standard sieve.

In another embodiment,

5

20

•25

the hydroxypropylmethyl cellulose of component b) has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the hydroxypropylmethyl cellulose of component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

- 30 In another embodiment, the solid dosage form comprises the following components:
 - a) a uniform admixture of :
 - (i) 500 mg/solid dosage form N-(2-Propylpentanoyl)glycinamide,

(ii) 50 mg/solid dosage form hydroxypropyl cellulose;and

- (iii) 100 mg/solid dosage form microcrystalline cellulose;
- b) 150 mg/solid dosage form of hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
 - c) 60 mg/solid dosage form of a different hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
 - d) 20 mg/solid dosage form lactose;
 - e) 4.5 mg/solid dosage form magnesium stearate; and
 - f) 1 mg/solid dosage form colloidal fumed silica.

20

30

35

15

In one embodiment, at least 90% of the hydroxypropylmethyl cellulose of component b), of component c), or of both component b) and c) passes through a No. 100 US standard sieve.

25 In another embodiment,

the hydroxypropylmethyl cellulose of component b) has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the hydroxypropylmethyl cellulose of component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

The subject invention also provides a hard compressed tablet comprising a uniform admixture of the following components:

- a) N-(2-Propylpentanoyl)glycinamide;
- b) a hydroxypropylmethyl cellulose; and
 - c) a different hydroxypropylmethyl cellulose.

In one embodiment,

5

10

15

20

25

30

the hydroxypropylmethyl cellulose component b) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve; and

the hydroxypropylmethyl cellulose component c) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

In one embodiment, at least 90% of the hydroxypropylmethyl cellulose of component b), of component c), or of both component b) and c) passes through a No. 100 US standard sieve.

In another embodiment,

the hydroxypropylmethyl cellulose component b) has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the hydroxypropylmethyl cellulose component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000

cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20° C.

In another embodiment, the tablet further comprises a filler, bubicant and flow agent as additional components.

In another embodiment,

10

20

25

the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant comprises sodium stearyl fumarate; and the flow agent comprises a colloidal fumed silica.

In another embodiment, the tablet comprises a uniform admixture of the following components:

- a) from 100 mg/tablet to 1000 mg/tablet N-(2-Propylpentanoyl)glycinamide;
- b) from 10 mg/tablet to 300 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;
- c) from 10 mg/tablet to 300 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;
- 30 d) from 1 mg/tablet to 300 mg/tablet a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
 - e) from 0.1 mg/tablet to 20 mg/tablet sodium steary1 fumarate; and
- f) from 0.1 mg/tablet to 15 mg/tablet a colloidal fumed silica.

In another embodiment, the tablet comprises a uniform admixture of the following components:

- a) from 400 mg/tablet to 1000 mg/tablet N-(2-Propylpentanoyl)glycinamide;
- b) from 100 mg/tablet to 300 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;
- c) from 20 mg/tablet to 150 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;
- d)from 10 mg/tablet to 60 mg/tablet a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
- e) from 2 mg/tablet to 20 mg/tablet sodium stearyl fumarate; and
- f) from 5 mg/tablet to 15 mg/tablet a colloidal fumed silica,
- 25 per 1 gram tablet.

10

15

20

In another embodiment, the tablet comprises a uniform admixture of the following components:

- a) 500 mg/tablet N-(2-Propylpentanoyl)glycinamide;
- b) 150 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds. (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;

c) 60 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;

d) 20 mg/tablet lactose;

5

15

25

- e) 10 mg/tablet sodium stearyl fumarate; and
- f) 10 mg/tablet colloidal fumed silica.
- 10 The subject invention also provides a composition in granulate form comprising a uniform admixture of:
 - (i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\begin{array}{c|c}
O & R_1 \\
N & (CH_2)n
\end{array}$$

$$NR_2R_3$$

or

O
$$R_1$$
 O NR_2R_3

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer

which is greater than or equal to 0 and less than or equal to 3; and

(ii) a hydroxypropyl cellulose.

In one embodiment of the composition, the active ingredient comprises a compound having the structure:

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

10 or

$$N$$
 (CH_2)
 N
 NR_2R_3

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3.

In another embodiment, the active ingredient comprises valproic 20 sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium or valpromide.

The subject invention also provides a tablet comprising the above granulate as a component.

25

15

In one embodiment of the tablet, the granulate further comprises a filler.

In another embodiment, the tablet further comprises a 5 hydroxypropylmethyl cellulose as a component.

In another embodiment, the tablet further comprises as additional components a filler, a lubricant and a flow agent.

10 In another embodiment, the tablet further comprises as additional components a filler, a lubricant and a flow agent.

In another embodiment, the tablet further comprises a different hydroxypropylmethyl cellulose as a component.

15

20

30

In another embodiment,

the hydroxypropylmethyl cellulose has 19%-24% by weight substituent, by 78-128 methoxyl hydroxylproproxyl substituent and has a particle size of the · 1east 99% that at distribution such hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

In another embodiment, at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

In another embodiment,

the hydroxypropylmethyl cellulose has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C.

In another embodiment, .

the different hydroxypropylmethyl cellulose has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

In another embodiment, at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

10

5

In another embodiment,

the different hydroxypropylmethyl cellulose has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

In another embodiment, the filler in the granulate is a microcrystalline cellulose.

20

15

In another embodiment,

the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

25 the lubricant comprises magnesium stearate, sodium stearyl fumarate or a combination thereof; and the flow agent comprises a colloidal fumed silica.

The subject invention also provides a sustained release tablet.

30 comprising a compound having the structure:

PCT/US2004/003281 WO 2004/071421

$$\bigcap_{N \in CH_2 \setminus n} \bigcap_{N \in R_2 \setminus R_3} \bigcap_{N \in R_3 \setminus R_3} \bigcap_{N \in R_3 \setminus R_3} \bigcap_{N \in R_3 \setminus R_3 \setminus R_3} \bigcap_{N \in R_3 \setminus R_3 \setminus R_3} \bigcap_{N \in R_3 \setminus R_3 \setminus R_3 \setminus R_3} \bigcap_{N \in R_3 \setminus R_3$$

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

5

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3.

10

 $N - \{2$ is compound embodiment, the one propylpentanoyl)glycinamide.

15 The subject invention also provides a method of treating neuropathic pain in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of any of the solid dosage forms or tablets of the invention in order to thereby treat the neuropathic pain in 20 the subject. .

The subject invention also provides a method of treating a headache disorder in a subject in need of such treatment comprising administering to the subject a therapeutically 25 effective dose of any of the solid dosage forms or tablets of

the invention in order to thereby treat the headache disorder in the subject.

The subject invention also provides a method of treating 5 epilepsy in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of any of the solid dosage forms or tablets of the invention in order to thereby treat epilepsy in the subject.

The subject invention also provides a method of controlling seizures in a subject suffering from epilepsy comprising administering to the subject a therapeutically effective dose of any of the solid dosage forms or tablets of the invention in order to thereby control the seizures in the subject.

15

The subject invention also provides a method of treating pain in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of any of the solid dosage forms or tablets of the invention in order to thereby treat pain in the subject.

The subject invention also provides a method of pain prophylaxis in a subject in need of such treatment comprising administering to the subject a prophylactic dose of any of the solid dosage forms or tablets of the invention in order to thereby effect pain prophylaxis in the subject.

The subject invention also provides a method of treating mania in bipolar disorder in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of any of the solid dosage forms or tablets of the invention in order to thereby treat mania in bipolar disorder in the subject.

The subject invention also provides a method of attenuating bipolar mood swings in a subject suffering from bipolar disorder comprising administering to the subject a therapeutically effective dose of any of the solid dosage forms or tablets of the invention in order to thereby attenuate the bipolar mood swings in the subject.

The subject invention also provides a process for preparing the above solid dosage form, comprising the steps of:

a) admixing predetermined amounts of

(i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

N $(CH_2)n$ NR_2R_3

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

20

25

10

15

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

-29-

- (ii) a binder;
- b) admixing the uniform mixture of step a) with a predetermined amount of a hydroxypropylmethyl cellulose; and
- 5 c) compressing the mixture of step b) to form the tablet.

In one embodiment of the process, step b) further comprises admixing the uniform mixture with a predetermined amount of a different hydroxypropylmethyl cellulose.

10

In another embodiment, step b) further comprises admixing the uniform mixture with predetermined amounts of a filler, a lubricant and a flow agent.

In another embodiment, the flow agent comprises colloidal fumed silica.

In another embodiment, the filler comprises microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing.

In another embodiment, the filler comprises lactose.

In another embodiment, the lubricant comprises magnesium stearate or sodium stearyl fumarate or a combination thereof.

In another embodiment, the lubricant comprises magnesium stearate.

30 In another embodiment,

each hydroxypropylmethyl cellulose of step b) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the

hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

In another embodiment, at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

In another embodiment,

the first hydroxypropylmethyl cellulose has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the second hydroxypropylmethyl cellulose has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

The subject invention also provides a process for preparing the 20 above hard compressed tablet comprising the steps of:

- a) admixing predetermined amounts of N-(2-Propylpentanoyl)glycinamide, hydroxypropylmethyl cellulose, and a different hydroxypropylmethyl cellulose; and
- b) compressing the mixture of step a) to form the hard compressed tablet.

In one embodiment,

each hydroxypropylmethyl cellulose of step a) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

10

In another embodiment, at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

In another embodiment,

the hydroxypropylmethyl cellulose has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the different hydroxypropylmethyl cellulose has an apparent viscosity of 6.138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

In another embodiment, step a) further comprises admixing predetermined amounts of a filler, lubricant and flow agent as additional components.

·20 In another embodiment,

15

the filler comprises microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant comprises sodium stearyl fumarate; and the flow agent comprises colloidal fumed silica.

The subject invention also provides a process for preparing the above composition in granulate form, comprising granulating a predetermined amount of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide or a compound having the structure:

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

5

10

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, and a predetermined amount of hydroxypropyl cellulose to form the composition in granulate form.

15 The subject invention also provides a process for preparing a sustained release tablet comprising the steps of:

- a) admixing the above granules with predetermined amounts of a hydroxypropylmethyl cellulose; and
- b) compressing the mixture of step a) to form the tablet.

20

In another embodiment, step a) further comprises admixing the granules with a predetermined amount of each of a different hydroxypropylmethyl cellulose, a filler, a lubricant and a flow agent.

In another embodiment, the flow agent comprises colloidal fumed silica.

In another embodiment, the filler comprises microcrystalline 5 cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing.

In another embodiment, the filler is lactose.

10 In another embodiment, the lubricant comprises magnesium stearate or sodium stearyl fumarate or a combination thereof.

In another embodiment, the lubricant comprises magnesium stearate.

15

20

.25

In another embodiment, the process comprises the steps of:

a) admixing the granules with predetermined amounts of hydroxypropyl methyl cellulose having an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C, and hydroxypropyl methyl cellulose having an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and b) compressing the mixture of step a) to form the tablet.

In another embodiment, step a) further comprises admixing the 30 granules with predetermined amounts of a flow agent, a filler, and a lubricant.

In another embodiment, the process comprises the steps of a) admixing the granules with

a predetermined amount of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C which results in tablets containing 150 mg/tablet;

a predetermined amount of hydroxypropyl methyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°Cwhich results in tablets containing 60 mg/tablet;

a predetermined amount of lactose which results in tablets containing 20 mg/tablet;

a predetermined amount of magnesium stearate which results in tablets containing 4.5 mg/tablet; and

a predetermined amount of a colloidal fumed silica which results in tablets containing 1 mg/tablet; and

b) compressing the mixture of step a) to form the tablet.

The subject invention also provides the use of an active 25 ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{\mathbf{N}} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

30

5

10

15

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form or tablet of the invention for use in treating a headache disorder in a subject.

The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)_{\Pi}} \bigcap_{NR_2R_3}$$

20

5

10

$$\bigcap_{\mathbf{N}} \bigcap_{(CH_2)\mathbf{n}} \bigcap_{\mathbf{N}R_2R_3}$$

or

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form or tablet of the invention for use in treating neuropathic pain in a subject.

10 The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$N$$
 $(CH_2)n$
 NR_2R_3

$$N$$
 $(CH_2)n$
 NR_2R_3

or

20

25

15

5

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group; or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage from or

-37-

tablet of the invention for use in treating epilepsy in a subject.

The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\begin{array}{c|c}
O & R_1 & O \\
N & (CH_2)n & NR_2R_3
\end{array}$$

10

15

20

or

$$\bigcap_{H}^{N} (CH_2)n \bigcap_{NR_2R_3}^{N}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid release dosage form or tablet of the invention for use in controlling seizures in a subject suffering from epilepsy.

The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of

valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

5 or

10

15

$$\bigcap_{N \to \infty} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form or tablet of the invention for use in treating mania in bipolar disorder in a subject.

The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$N$$
 $(CH_2)n$
 NR_2R_3

or

$$\begin{array}{c|c}
O & R_1 & O \\
N & (CH_2)n & NR_2R_3
\end{array}$$

5

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form or tablet of the invention for use in attenuating bipolar mood swings in a subject suffering from bipolar mood disorder.

10

The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\begin{array}{c|c}
O & R_1 & O \\
N & (CH_2)n & NR_2R_3
\end{array}$$

20

$$\bigcap_{H}^{N} \bigcap_{(CH_2)n}^{(CH_2)n} \bigcap_{NR_2R_3}^{N}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form or tablet of the invention for use in treating pain in a subject.

10

5

The subject invention also provides the use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

or

$$\bigcap_{N \to \infty} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

20

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form or tablet of the invention for use in effecting pain prophylaxis in a subject.

The subject invention also provides the sustained release solid dosage form or tablet for use in treating a headache disorder in a subject.

The subject invention also provides the sustained release solid dosage form or tablet for use in treating neuropathic pain in a subject.

The subject invention also provides the sustained release solid dosage form or tablet for use in treating epilepsy in a subject.

20

5

The subject invention also provides the sustained release solid dosage form or tablet for use in controlling seizures in a subject suffering from epilepsy.

25 The subject invention also provides the sustained release solid dosage form or tablet for use in treating mania in bipolar disorder in a subject.

The subject invention also provides the sustained release solid 30 dosage form or tablet for use in attenuating bipolar mood swings in a subject suffering from bipolar disorder.

The subject invention also provides the sustained release solid dosage form or tablet for use in treating pain in a subject.

35

The subject invention also provides the sustained release solid dosage form or tablet for use in effecting pain prophylaxis in a subject.

The subject invention also provides a controlled release oral unit dose composition comprising N-(2-propylpentanoy1) glycinamide and at least one pharmaceutically acceptable carrier, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoy1) glycinamide between 4 and 24 hours after ingestion of a single oral unit dose.

In one embodiment, the composition when orally ingested by a 10 human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 4 and 12 hours after ingestion of a single oral unit dose.

In a further embodiment, the composition, when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 6 and 12 hours after ingestion of a single oral unit dose.

In a further embodiment, the composition, when orally ingested 20 by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 6 and 8 hours after ingestion of a single oral unit dose.

In a further embodiment of the above controlled release oral dose compositions, the peak blood plasma level of N-(2-propylpentanoyl) glycinamide is from 0.5 micrograms/ml to 16 micrograms/ml per a 1000 mg dose of N-(2-propylpentanoyl) glycinamide in the composition.

30 In another embodiment, the composition, when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycine in the human subject from 0.5 μg/mL to 1.7 μg/mL per a 1000 mg dose of N-(2-propylpentanoyl) glycinamide in the composition.

.35

The subject invention also provides a controlled release oral dose composition comprising N-(2-propylpentanoyl) glycinamide

5

and a pharmaceutically acceptable carrier, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide of 0.5 µg/mL to 16 µg/mL per a 1000 mg dose in the composition.

The subject invention also provides a controlled release oral dose composition comprising N-(2-propylpentanoy1) glycinamide and a pharmaceutically acceptable carrier, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoy1) glycine of 0.5 μ g/mL to 1.7 μ g/mL per a 1000 mg dose of N-(2-propylpentanoy1) glycinamide in the composition.

The subject invention also provides a method of inducing in a 15 human subject peak blood plasma level of N-(2propylpentanoyl) glycinamide between 4 and 24 hours after administration of N-(2-propylpentanoyl) glycinamide, comprising administering to the human subject a controlled release oral unit dose composition comprising N-(2-propylpentanoy1) 20 glycinamide and at least one pharmaceutically acceptable carrier, which composition induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 4 and 24 hours after administration of a single oral unit dose.

25 In one embodiment, the peak blood plasma Level of N-(2-propylpentanoyl) glycinamide occurs between 4 and 12 hours after administration.

In another embodiment, the peak blood plasma level of N-(2-30 propylpentanoyl) glycinamide is 0.5 µg/mL to 16 µg/mL per 1000 mg dose of N-(2-propylpentanoyl) glycinamide in the composition.

In one embodiment of any of the above methods, the

35 administeration to the human subject of a controlled release

oral unit dose composition comprising N-(2-propylpentanoy1)

glycinamide and at least one pharmaceutically acceptable carrier induces a peak blood plasma level of N-(2-propylpentanoy1) glycine in the human subject from 0.5 µg/mL to 1.7 µg/mL upon administration of a single 1000 mg dose of N-(2-propylpentanoy1) glycinamide.

In another embodiment, the controlled release oral dose composition is any of the solid dosage forms or the tablets described above.

10

In another embodiment of the invention, the process for manufacturing the sustained release formulation of N-(2-Propylpentanoyl)glycinamide comprises:

- 1. Preparing a granulate of N-(2-Propylpentanoyl)glycinamide
- 2. Mixing the granulate of step 1 with excipients
 - 3. Compressing the mixture of step 2 to form a sustained release tablet of N-(2-Propylpentanoyl)glycinamide

In another embodiment, the process for manufacturing the sustained release formulation of N-(2-Propylpentanoyl)glycinamide comprises:

- 1. Mixing the active material with a carrier and other excipients
- 2. Direct compression of the mixture of step 1.

25

30

In another embodiment, the process for manufacturing the sustained release formulation of N-(2-Propylpentanoyl)glycinamide comprised:

- Mixing N-(2-Propylpentanoyl)glycinamide with a carrier and other excipients
 - 2. Compression of the mixture of step 1 into tablets
 - 3. Preparing slugs of the tablets of step 2
 - 4. Filling into capsules the slugs of step 3

In another embodiment, the process for manufacturing the sustained release formulation of N-(2-propylpentanoyl)glycinamide comprised:

1. Mixing N-(2-Propylpentanoyl)glycinamide with a carrier and other excipients

- 2. Compression of the mixture of step 1 into tablets
- 3. Preparing slugs of the tablets of step 2
- 4. Dispersing the slugs of step 3 in suspension

"Slugs" are granulates manufactured via a dry granulation process that involves milling the tablets into small particles.

10 The present invention provides a sustained release pharmaceutical composition comprising the active-material N-(2-Propylpentanoyl)glycinamide.

The subject invention also provides an oral dosage of N-(2-15 Propylpentanoyl)glycinamide sustained release form.

As used herein, "US Standard Sieve No. 40" refers to a sieve having a specified sieve opening of 0.0165 inches and a specified wire diameter of 0.0098 inches.

20

release" does not.

As used herein, "US Standard Sieve No. 100" refers to a sieve having a specified sieve opening of 0.0059 inches and a specified wire diameter of 0.0040 inches.

- As used herein, the phrase "controlled release" dosage forms refer to dosage forms which are formulated to release the drug slowly over a prolonged period of time. These dosage forms are also referred to as "sustained-release" or "prolonged release" dosage forms (Remington: The Science and Practice of Pharmacy, 20th ed. P. 859). However, the term "controlled release" also includes enterically coated tablets while the term "sustained
- As used herein, the term "compressed tablets" refers to tablets which formed by a press tableting machine which applies a

compression force of between about 2000 lb (about 8.9 \times 10^3 Newtons) and about 10,000 lb (4.45 \times 10^4 Newtons).

As used herein, the term "hard compressed tablets" refers to tablets which remain unchanged under compression forces ranging from about 2000 lb (1.3 x 10⁴ Newtons) to about 10,000 lb (4.45 x 10⁴ Newtons). The term "hard compressed tablets" does not include within its scope any granulate which does not itself meet the test for hardness described above.

10

There are several in vitro mechanisms by which the N-(2-Propylpentanoyl)glycinamide can be released. One such mechanism is sustained release in matrix tablets. The main principle of this mechanism is that the water partially hydrates the outer layers of the tablet to form a gel layer. Throughout the life of the ingested tablet, the rate of drug diffusion and of the wet gel and the rate of the tablet erosion control the overall dissolution rate and drug availability.

20 This matrix can be obtained by direct compression or by initial granulation, which granules are then compressed into the matrix system. In monolithic matrix systems, the drug is homogeneously dispersed throughout a polymer mass of other carrier material.

25 Release characteristics depend on the geometry of the system, the nature of the polymer and other excipients, solubility and the processing methods.

N-(2-Propylpentanoyl)glycinamide is difficult to work with due to its "lamination and compression" characteristics. To alleviate the problem of lamination, the subject invention employs a filler and hydroxypropylmethyl cellose as a carrier which improve the compressing characteristics while simultaneously slowing down the release profile.

In a preferred embodiment, the carrier is Methocel kl00 LV, and the filler is lactose.

As described more fully in the examples which follow, in order to develop a prototype with a slower dissolution profile, the concentration of the carrier (e.g. methocel) was increased until any further increase gave no effect on the resulting dissolution profile. At this point, the polymer had achieved the maximum sustained action.

10

In order to further improve the dissolution profile, a second molecular weight grade of methocel was added to the formulation. While the first grade of Methocel improved the compression properties and achieved a maximum sustained action, the second grade detracted from the physical characteristics of the tablet but improved the sustained-release action. However, by combining these two different molecular weight grades of methocel in the correct proportions, the dissolution rate was decreased and the tablets were made with the desired physical characteristics.

Thus, the subject invention provides a sustained release formulation of N-(2-Propylpentanoyl)glycinamide which contains two different grades of Methocel combined in the correct proportions to achieve the desired dissolution profile and the desired compressibility characteristics.

In the case of compression tablets, the excipients give the desired flow of granules, and uniform compressibility into 30 tablets.

The pharmaceutical excipients include fillers, flow agents, disintegrants and lubricants.

Most multiparticulate systems are delivered in the form of solid dosage. However, for some patients, it is desirable to use extended release dosage forms in liquid form. The multiparticulate system can be a redispersable dosage form, or a liquid suspension.

Non-limiting examples of a filler used in the subject invention (used for example for weight adjustment and for better compression) are corn starch, lactose, glucose, various natural gums, methylcellulose, carboxymethylcellulose, microcrystalline cellulose (e.g. Avicel® PH101 or 102 (FMC Corporation, Philadelphia, PA)), calcium phosphate, calcium carbonate, calcium sulfate kaolin, sodium chloride, powdered cellulose, sucrose, mannitol and starch. In a preferred embodiment, the excipient useful as a filler comprises a microcrystalline cellulose.

Non-limiting examples of a carrier (extended release agent) used in the subject invention (used for example for the 20 controlled release) are cellulose acetate, glycery1 monostearate, zein, microcrystalline wax, hydroxymethylcellulose, hydroxyethylcellulose, hydroxypropylmethylcellulose, hydroxypropylcellulose Klucel®), carboxyvinyl polymers, polyvinyl alcohols, glucans, 25 scleroglucans, chitosans, mannans, galactomannans, alginic acid and salts and derivatives thereof, acrylates, methacrylates, acrylic/methacrylic copolymers, polyanhydrides, polyaminoacids, methyl vinyl ethers/maleic anhydride copolymers, carboxymethylcellulose and derivatives thereof, 30 ethylcellulose, methylcellulose and cellulose derivatives in general, modified starch and polyesters, polyethylene oxide.

In an embodiment, the excipient used as a carrier comprises a hydroxypropylmethylcellulose. In another embodiment, the hydroxypropylmethylcellulose has an average molecular weight

between about 10 kDa and about 1500 kDa. In a further embodiment, the hydroxypropylmethylcellulose has methoxyl substituent and 7%-12% hydroxylproproxyl substituent. In an added embodiment, the hydroxypropylmethyl cellulose has a 5 pH of 5.5-8.0 in a 1% solution. In an added embodiment, the hydroxypropylmethylcellulose has a particle size distribution such that about 100% of the hydroxypropylmethylcellulose passes through 30 mesh screen. In one embodiment, hydroxypropylmethylcellulose has a particle size distribution 10 such that about 99% of the hydroxypropylmethylcellulose passes through a 40 mesh screen. In yet another embodiment, the hydroxypropylmethylcellulose has a particle size distribution such that 55%-95% of the hydroxypropylmethylcellulose passes through a 100 mesh screen. In yet another embodiment, the 15 hydroxypropylmethylcellulose has a particle size distribution such that 90% of the hydroxypropylmethylcellulose passes through a 100 mesh screen. In a further embodiment, the hydroxypropylmethylcellulose has a particle size distribution such that 65%-85% of the hydroxypropylmethylcellulose passes 20 through a 100 mesh screen. In an additional embodiment, the hydroxypropylmethylcellulose has a particle size distribution such that about 80% of the hydroxypropylmethylcellulose passes through a 100 mesh screen. In a further embodiment, the hydroxypropylmethylcellulose has a particle size distribution 25 such that about 90% of the hydroxypropylmethylcellulose passes through a 100 mesh screen. In a further embodiment, the hydroxypropylmethylcellulose is a Methocel® polymer (Colorcon, West Point, PA), such as Methocel® K100 Premium LV EP or LV LH EP alone or in combination, or Methocel® K15M EP or CR EP.

30

Non-limiting examples of a binding agent used in the subject invention (used for example for the granulate) are alginic acid, acia, carbomer, carboxymethylcellulose sodium, dextrin, ethylcellulose, gelatin, guar gum, hydrogenated vegetable oil, hydroxyethylcellulose, hydroxypropylcellulose (e.g., Klucel®,

Aqualon Division, Hercules Incorporated, Wilmington, Del.), glucose, magnesium hydroxypropylmethylcellulose, liquid methylcellulose, maldodextrin, aluminum silicate, polymethacrylates, povidone, pregelatinized starch, 5 alginate, starch, and zein. In a preferred embodiment, the agent comprises excipient used as a binding hydroxypropylcellulose.

embodiment, the excipient used as a binder is In 10 hydroxypropyl cellulose. In one embodiment, the hydroxypropyl cellulose has a particle size distribution such that about 85% of the hydroxypropyl cellulose passes through a 30 mesh screen. In another embodiment, the hydroxypropyl cellulose has a particle size distribution such that about 99% of 15 hydroxypropyl cellulose passes through a 20 mesh screen. In another embodiment, the hydroxypropyl cellulosehas a pH of 5.0-7.5 in water solution. In one embodiment, the hydroxypropyl cellulose has an average molecular weight of 1,150,000. In one embodiment, the hydroxypropyl cellulose has an 20 molecular weight of 850,000. embodiment, the In one hydroxypropyl cellulose has an average molecular weight of 370,000. In one embodiment, the hydroxypropyl cellulose has an average molecular weight of 140,000. In one embodiment, the hydroxypropyl cellulose has an average molecular weight of 25 95,000. In one embodiment, the hydroxypropyl cellulose has an average molecular weight of 80,000. In one embodiment, the hydroxypropyl cellulose has a viscosity of 1,500-3,000 cps at a concentration of 1% by weight in water at 25°C. embodiment, the hydroxypropyl cellulose has a viscosity of 30 4,000-6,500 cps at a concentration of 2% by weight in water at 25°C. In one embodiment, the hydroxypropyl cellulose has a viscosity of 150-400 cps at a concentration of 2% by weight in water at 25°C. In one embodiment, the hydroxypropyl cellulose has a viscosity of 150-400 cps at a concentration of 5% by 35 weight in water at 25°C. In one embodiment, the hydroxypropy1

cellulose has a viscosity of 75-150 cps at a concentration of 5% by weight in water at 25°C. In one embodiment, the hydroxypropyl cellulose has a viscosity of 200-600 cps at a concentration of 10% by weight in water at 25°C. In one embodiment, the hydroxypropyl cellulose has a viscosity of 75-150 cps at a concentration of 5% by weight in water at 25°C. In one embodiment, the hydroxypropyl cellulose has a viscosity of 300-600 cps at a concentration of 10% by weight in water at 25°C.

10

In one embodiment, the excipient used as a filler is a microcrystalline cellulose. In an added embodiment, the microcrystalline cellulose has an average particle size between about 50 and about 90 microns.

15

Non-limiting examples of a flow agent used in the subject invention are micron-sized silica powders. A non-limiting example of a flow agent used in the subject invention (used for better flow of the mix for compression) is colloidal silicon dioxide or Syloid®.

Non-limiting examples of a lubricant used in the subject invention (used for example for better compression properties) are talc, sodium stearyl fumarate, magnesium stearate, calcium stearate, hydrogenated castor oil, hydrogenated soybean oil and polyethylene glycol (PEG) or combinations thereof.

Details of general formulation procedures and information on additional excipients may be found in Remington: The Science and Practice of Pharmacy, 20th Edition.

This invention will be better understood from the Experimental Details which follow.

Experimental details

Example 1: Manufacture of N-(2-Propylpentancyl)glycinamide Sustained Release (SR) tablets:

Granules of N-(2-Propylpentanoyl)glycinamide:

N-(2-n-Propylpentanoyl) glycinamide was granulated with a binder solution and with several excipients.

Table 1: Composition of the granules

Excipient	Use	Mg/tablet
N-(2-Propylpentanoyl) glycinamide	Active material	500
Microcrystalline Cellulose	Filler	100
Hydroxypropyl cellulose	Binder	50
Total		650

The tablets were then prepared by mixing the granulate with a carrier/carriers and several excipients (table 2).

15

10

Table 2: Composition of the sustained release tablets

		Α	. B	С
Excipient	Use		Mg/Tablet	
N-(2-Propylpentanoyl) glycinamide Granulate		650	650	, 650
Aerosil	Flow-agent	1.0	1.0	
Lactose	Filler	80	20	145
Hydroxypropyl Methyl Cellulose (Methocel K15M)	Carrier		60	
Hydroxypropyl Methyl Cellulose (Methocel K 100LV)	Carrier	150	150	
Magnesium Stearate	Lubricant	4.5	4.5	6
Crosscarmelose Sodium	Disintegrant			50

Each formulation was tested in a dissolution test using 900 ml purified water, 37°C, in US Pharmacopoeia (USP), versus the immediate release formulation (Formulation C).

Table 3: Dissolution of N-(2-Propylpentanoyl)glycinamide SR tablets

Formula	A	В	С	
Time (h)	% Dissolution			
0.5	7	4	100	
2	34	15		
4	66	32		
6	88	48		
10	102	75		
12		86		
14		96		
16		102		

As can be seen two different prototypes (A, B) of N-(2-5 Propylpentanoyl)glycinamide sustained release characteristics were observed.

Example 2: Effect of carrier on dissolution rate

10 Each of the following formulations contained different carriers in order to determine the effect of the carrier on the dissolution rate.

Table 4: Variations in the carriers

Formula	1	D	E	F	G	Н
					Methocel K15M*	
Excipient	Use			Mg/Tablet		
N-(2- Propylpentanoyl) glycinamide Granulate		650	650	650	650	650
Aerosil	Flow- agent	16.5	16.5	16.5	16.5	16.5
Lactose	Filler	80	80	80	80	80
*Carrier	*Carrier	120	120	120	100	120
Magnesium Stearate	Lubricant	4.5	4.5	4.5	4.5	4.5

15

Each formulation was tested in a dissolution test using 900 ml purified water 37°C, in US Pharmacopoeia (USP). The dissolution profile was found to be dependent upon the type of the carrier.

Table 5: Dissolution of tablets D-H

Formula	D Methocel K100M	E Klucel HF	F Carbopol 974p	G Methocel K100LV	H Methocel K15M
Time (h)	,		% Dissolution		
0.5	2	5	7	12	7
1	4	8	16	26	. 15
2 .	8	12	26	52	32
3	10	16	. 30	71	47
4	13	19	33	86	61
6	17	25	39	106	83
8	21	30	45		100
12	29	40	54		

Due to their resulting dissolution profile of 6-8 hours, 5 Methocel K100LV and/or Methocel K15M were selected as suitable carriers.

Example 3: Effect of the amount of carrier on dissolution rate

In order to determine the effect of the amount of the carrier on the dissolution rate, formulations were tested while varying the amount of Methocel K100 LV and/or Methocel K15M.

Table 6: Variation in the amount of the carrier (Methocel K100 LV)

Form	ula	1	J	K
Excipient	Use		Mg/Tablet	
N-(2- Propylpentanoyl) glycinamide Granulate		650	650	650
Aerosil	Flow-agent	1.0	1.0	1.0
Lactose	Filler	80	80	. 60
Hydroxypropyl Methyl Cellulose (Methocel K100 LV)	Carrier	100	150	170
Magnesium Stearate	Lubricant	4.5	4.5	4.5

Each formulation was then tested in a dissolution test using 900 ml purified water, 37°C, in US Pharmacopoeia (USP).

15

Table 7: Dissolution of formulations I-K

Table 7. Dissolution of formalations 1-10					
Formula	l	J	K		
Time (h)	9	% Dissolution			
0.5	15	11	8		
1	28	20	12		
2	49	39	35		
3	64	54	51		
4	76	68	65		
6	94	87	87		
8	104	98	102		
12	105	105	110		

The results showed that the dissolution profile was dependent upon the amount of the carrier (Methocel K100LV). Increasing the concentration of the polymer in the matrix system increases the viscosity of the outer layer gel which forms and leads to a more delayed release of the drug product. However, when increasing the amount of carrier from formulation J to formulation K the effect on the endpoint of the dissolution was less significant than the change observed when changing from formulation I to J. Thus, formulation J achieves the maximum sustained action for this polymer.

The same procedure was followed in order to determine the effect of the amount of Methocel® K15M on the dissolution profile.

Table 8: Variation in the amount of the carrier (Methocel K15M)

Formi	ula	L	M	N
Excipient	Use		Mg/Tablet	
N-(2- Propylpentanoyl) glycinamide Granulate	·	650	650	650
Aerosil	Flow-agent	1.0	1.0	1.0
Lactose	Filler	80	80	60
Hydroxypropyl Methyl Cellulose (Methocel K15M)	Carrier	80	100	150
Magnesium Stearate	Lubricant	4.5	4.5	4.5

Each formulation was tested in a dissolution test using 900 ml purified water, 37°C, in US Pharmacopoeia (USP).

Table 9: Dissolution of formulations L-N

Formula	L	M	N
Time(h)	•	% Dissolution	າ
0.5	9	6	4
1	13	10	7
2	21	17	12
3	29	23	17
4	37	30	22
6	50	43	31
8	63	55	40
10	75	66	49
12	84	75	57
14		84	
16		. 90	72
18			76

The results showed that the dissolution profile was dependent upon the amount of the carrier (Methocel® K15M) and that increasing the concentration of this polymer in the matrix system delays the release of the drug product. More polymer in the matrix leads to more polymer on the tablet surface. Hence, wetting is more easily achieved and gel formation is accelerated. However, formulation N suffered from poor compressibility characteristics.

The use of Methocel K15M as a carrier was found to slow the dissolution profile. However, it also yielded tablets with poor compressibility properties. Other alternatives were therefore investigated in order to produce tablets with good compressibility properties as well as slow dissolution profiles.

Example 4: Effect of time from production on dissolution rate

Table 10: Time effect of production on the dissolution profile

		0	P
Excipient	Use	Mg/T	Tablet
N-(2- Propylpentanoyl) glycinamide Granulate		650	650
Aerosil	Flow-agent	1.0	1.0
Lactose	Filler	80	80
Hydroxypropyl Methyl Cellulose (Methocel K100 LV)	Carrier	150	150
Magnesium Stearate	Lubricant	4.5	4.5

O: The tablets were kept in uncontrolled conditions for two years.

The formulations were then checked for dissolution profile.

Table 11: Dissolution of tablets O-P

Formula	0	Р
Time(h)_	% Dis	solution
0.5	9	11
1	20	20
2	39	39
3	55	54
4	67	68
6	· 85	87
8	95	98
12	101	105

10

The results indicated that the formulations utilized were extremely stable.

⁵ P: The same formulation was compressed anew.

Example 5: Effect of combining Methocel carriers on dissolution rate

Table 12: Effect of combining different amounts of of Methocel carriers.

Formula		Q	S
Excipient	Use	iviy/Tablet	
N-(2- Propylpentanoyl) glycinamide Granulate		650	650
Aerosil	Flow- agent	1.0	1.0
Lactose	Filler	40	40
Carrier (Methocel K100LV)	Carrier	150	150
Carrier (Methocel K15M)	Carrier	40	75
Magnesium Stearate	Lubricant	4.5	4.5

Each formulation was tested in a dissolution test using 900 ml purified water, 37°C, in US Pharmacopoeia (USP).

Table 13: Dissolution profile of tablets Q and S

Formula	Q	S
Time (h)	% Diss	olution
0.5	6	5
2	23	18
4	46	34
6	65	49
8	80	
10	92	73
12	100	87
14	105	91
16		94

As illustrated by the above results, increasing the amount of Methocel K15M relative to Methocel K100 LV improved and decreased the dissolution rate.

Example 6: Effect of lubricant type and amount on dissolution rate

Table 14: Effect of Lubricant type and amount

Formula	ν	W	Χ	Υ	
Excipient	Use		Mg/ĭ	abiel	
N-(2-n- Propylpentanoyl) glycinamide Granulate		650	650	650	650
Aerosil	Flow-agent	1.0	1.0	1.0	1.0
Lactose	Filler	20	20	80	80
Methocel K100LV	Carrier	150	150	150	150
Methocel K15M	Carrier	60	60		
Magnesium Stearate	Lubricant	4.5		4.5	
Sodium Stearyl Fumarate (Pruv)	Lubricant		9.0		9.0

Each formulation was tested in a dissolution test using 900 ml purified water, 37°C, in US Pharmacopoeia (USP).

Table 15: Dissolution profile of tablets V-Y

Formula	V	W	X	Υ
Time (h)		% Dis	solution	-
0.5	4	4	7	7
2	15	16	34	34
4	32	34	66	66
6	48	52	88	89
10	75	78	102	103
12	86	88		
14	96	95		
16	102	101		
18		105		

As can be seen, no effect on dissolution profile was observed when changing the lubricant type or quantity. However, the physical compressing properties were improved when using Pruv

instead of magnesium stearate.

10

5

Example 7: Effect of apparatus type on dissolution rate

Table 16: Influence of apparatus type

Formulation	A	В	
Excipient	Use .	mg/tablet	mg/tablet
N-(2-	Active	650	650
propylpentanoyl)glycinamide			j
granulate			
Aerosil	Flow-agent	1.0	1.0
Lactose	Filler	80	20
Methocel K100LV	Carrier	150	150
Methocel K15M	Carrier	0	60
Magnesium Stearate	Lubricant	4.5	4.5

The formulations were tested using dissolution tests using two different apparatuses using 900 mL purified water at 37°C, according to US Pharmacopoeia (USP). Apparatus 1 (basket apparatus) was maintained at 100 RPM. Apparatus 2 (paddle apparatus) was maintained at 75 RPM.

Table 17: Dissolution profile of tablets according to Apparatus 1 and Apparatus 2

	Formul	ation A	Formula	ation B
	Apparatus 1	Apparatus 2	Apparatus 1	Apparatus 2
Time	% diss	olution	% diss	olution
(h)				
0.5	7	8	4	4
2	. 34	34	15	18
4	66	64	32	36
6	88	. 87	48	53
10	108	104	75	79
12			86	88
14			96	95
16			102	105

Results: The apparatus type used did not significantly influence the dissolution rate.

Example 8: Effect of manufacturing procedure on dissolution 5 rate

Table 18: Direct compression (DC, DC1, DC2, DC3) versus wet granulation (W)

Formula	W	DC	DC1	DC2	DC3	
Excipient	Use.		mg	/tablet		-
N- (2-	Active	650	500	750	750	750
propylpentanoyl)		granulate	active	active	active	active
glycinamide			only	only	only	only
Aerosil (syloid)	Flow	1.0	10	15	15	15
	agent					
Lactose	Filler	20	20			
Methocel K100LV	Carrier	150	150	150	150	150
Methocel K15M	Carrier	60	60			
Methocel K15MCR	Carrier			60	100	60
Pruv	Lubricant	. 9	10	15	15	20

10 The formulations were tested in standard dissolution tests using 900 ml purified water at 37°C according to USP.

Table 19: Dissolution profile of tablets

Formula	W	DC	DC1	DC2	DC3
Time (h)		% d:	ssolution		
0.5	4	7	13	8	12
2	16	18	27	18	27
4	34	32	44	31	43
6	52	44	58	42	59
10	78	67	. 71	- 53	72
12	88	77	83	63	85
14	95	85	94	73	96
16	101	91	100	81	101
18	105	95	102	88	103

These results show that although the active material is difficult to work with due to its unsatisfactory compression characteristics, direct compression technology and wet granulation technology both yielded tablets with a slow dissolution profile.

Example 9: Effect of amount of lactose on dissolution rate

Table 20: Effect of lactose on the dissolution rate

		EE	FF	GG	HH	- II	JJ
Exciplent	Use			Mg/T	ablet		
N-(2- Propylpentanoyl) glycinamide Granulate		650	650	650	650	650	650
Aerosil	Flow- agent	1.0	1.0	1.0	1.0	1.0	1.0
Lactose	Filler	40		40		80	40
Methocel K100LV	Carrier	150	150	150	150	100	100
Methocel K15M	Carrier	40	40	75	75		
Magnesium Stearate	Lubricant	4.5	4.5	4.5	4.5	4.5	4.5

10

Each formulation was tested in a dissolution test using 900 ml purified water, 37°C, in US Pharmacopoeia (USP).

Table 21: Dissolution profile of tablets EE-JJ

Formula	EE	FF	GG	НН	11	JJ				
Time (h)		% Dissolution								
0.5	6	5	5	4	13	11				
2	23	19	18	17	48	42				
4	46	38	34	34	82	71				
6	65	. 55	49	49	101	87				
8	80	69		65	106	96				
10	92	81	73	73	106	100				
12	100	91		90	107	100				
14	104	98	87	96		100				
16		101	91	100						

15

As can be seen, in formulations containing higher amounts of lactose, the dissolution rate was faster. However, the influence of the lactose decreased when the amount of the

carrier increased and the influence of the carrier became more effective.

Example 10: Additional testing on the effect of different lubricants and pH on dissolution profile

Table 22: Formulations tested

		Α	В	W	Υ
Excipient	Use		Mg/	Tablet	
N-(2-Propylpentanoyl) glycinamide Granulate		650	650	650	650
Aerosil	Flow- agent	1.0	1.0	1.0	1.0
Lactose	Filler	80	20	20	80
Hydroxypropyl Methyl Cellulose (Methocel K15M)	Carrier		60	60	
Hydroxypropyl Methyl Cellulose (Methocel K 100LV)	Carrier	150	150	150	150
Magnesium Stearate	Lubricant	4.5	4.5		
Pruv	Lubricant			9.0	9.0

10

Table 23: Dissolution profiles. Formula A with Magnesium Stearate

Intervals	Basket	Paddle	Basket	Paddle	Paddle	Basket
(min.)	100rpm Water	75rpm Water	100rpm 0.1N HCI	75rpm Intestinal fluid	75rpm Gastric fluid	100rpm Phosphate pH≃6,8
30	7	8	7	8	7	7
60	15	17	16	16	15	14
120	33	34	33	32	31	29
180	50	50	48	46	44	44
240	63	65	61	57	57	57
360	85	87	82	78	76	77
480	98	99	98	91	89	92
600	103		102			101
720	103		103			103

Table 24: Dissolution profiles. Formula B with Magnesium Stearate

Intervals	Basket	Paddle	Basket	Paddle	Paddle	Basket
(min.)	100rpm Water	75rpm Water	100rpm 0.1N HCI	75rpm Intestinal fluid	75rpm Gastric fluid	100rpm Phosphate pH=6.8
30	4	4	4	4	3	4
60		8		7	7	
120	15	16	15	14	13	13
180		24		20	19	
240	31	31	31	26	24	25
360	46	45	45	37	36	37
480	60	58	59	48	46	47
600	72		70			56
720	83		80			65
840	93		89			72
960	100		95			79

Table 25: Dissolution profiles. Formula Y

	Basket	Basket	
Intervals	100rpm	100rpm	
(min.)	Water	Gastric fluid	
30	7	6	
60		14	
120	34	29	
180			
240	66	57	
360	89	78	
480			
600	103		
720	104		

Table 26: Dissolution profiles. Formula W

	Basket	Basket	
Intervals	100rpm	100rpm	
(min.)	Water	Gastric fluid	
30	4	3	
60		7	
120	16	14	
240	34	29	
360	52	45	
600	78		
· 720	88		
840	95		
960	101		

10

5

As the results show, release rates of drug are unaffected by pH as the viscosity of the gel which forms on the tablet surface

and the rate of hydration are relatively independent of the pH environment. However, when ionic salts are used in the dissolution medium they can compete with the polymer and affect the dissolution rate of drug.

5

Example 11

Plasma Concentration of N-(2-propylpentanoyl) glycinamide and of N-(2-propylpentanoyl) glycine after administration.

10 Formulations A, B, and C were prepared as described in Example 1.

Two tablets of formulation A (2 X 500 mg active pharmaceutical ingredient) were simultaneously administered to each of 18 healthy male Caucasian volunteers. Plasma concentrations of N-(2-propylpentanoyl) glycinamide and of a major metabolite, N-(2-propylpentanoyl) glycine of each of the volunteers were regularly analyzed at 0.25, 0.5, 1.0, 1.5, 2, 4, 6, 8, 10, 12, 14, 16 and 24 hours.

20

The trial was then repeated with formulations B and C. The results of the trial were averaged and the mean plasma concentrations after administration of each of the formulations are depicted in figures 1 and 2.

25

Table 27 C_{max} and T_{max} of N-(2-propylpentanoyl) glycinamide after administering Formulations A, B, and C

Subject		C _{max} (µg/ml)			T _{max} (µg/ml)	
	Formulation	Formulation	Formulation	Formulation	Formulation	Formulation
	Α	В	С	. А	В	С
1	10.98	7.22	22.83	4.00	4.00	2.00
2	. 9.85	8.54	32.06	6.00	_ 16.00	0.25
3	9.77	7.39	22.67	10.00	16.00	2.00
4	9.39	7.91	19.48	12.00	16.00	0.50
5	10.12	6.07	Not Available	6.00	14.00	Not Available
6	12.89	7.64	34.33	4.00	4.00	0.50
7	10.85	7.61	29.96	12.00	6.00	1.00
8	9.65	7.44	21.26	6.00	16.00	0.50
9	10.95	8.03	20.39	6.00	14.00	1.50
10	8.32	6.48	21.52	6.00	6.00	1.00
11	9.18	6.78	17.19	14.00	14.00	4.00
12	8.86	8.53	21.28	6.00	14.00	0.50
13_	13.89	8.33	17.59	6.00	24.00	4.00
14	11.37	7.58	18.18	6.00	6.00	2.00
15	9.92	6.84	46.52	4.00	12.02	0.50
16	12.14	9.56	33.48	4.00	4.00	0.50
17	10.12	7.76	34.91	6.00	6.00	0.50
18	11.60	8.40	19.77	4.00	14.00	0.50
N	18	18	17	18	18	17
Mean	10.55	7.67	25.50	6.78	11.45	1.28
SD	1.44	0.84	8.22	3.08	5.73	1.19
Min	8.32	6.07	17.19	4.00	4.00	0.25
Median	10.12	7.63	21.52	6.00	14.00	0.50
Max	13.89	9.56	46.52	14.00	24.00	4.00

 C_{max} is the maximum measured plasma concentration of N-(2-5 propylpentanoyl) glycinamide after administration. T_{max} is the time at which the maximum concentration of N-(2-propylpentanoyl) glycinamide was measured.

As seen in figure 1, formulations A and B maintain a mean 10 plasma concentration of N-(2-propylpentanoyl) glycinamide which is stable from 4 hours after administration to 16 hours after administration. In addition, in formulations A and B, mean T_{max}

occurs after 6 hours, whereas in formulation C, mean T_{max} occurs before 2 hours.

As seen in table 27, the C_{max} after administration of formulations A and B did not exceed 14 µg/ml in any of the volunteers. However, the mean C_{max} after administration of formulation C was 25.5 µg/ml. Administration of formulations A or B may eliminate unwanted side-effects which are caused as a result of dosage peaks present in immediate release formulations such as formulation C.

Table 28 C_{max} and T_{max} of N-(2-propylpentanoy1) glycine after administering Formulations A, B, and C

	C _{max} (µg/ml)			T _{max} (µg/ml)			
Subject	Formulation	Formulation	Formulation	Formulation	Formulation	Formulation	
	A	B	C	A	В	С	
1	1.36	0.90	3.16	4.00	4.00	2.00	
2	1.27	1.27	2.68	4.00	23.88	0.50	
3	1.18	0.81	2.71	10.00	8.00	2.00	
4	1.11	0.99	2.24	12.00	16.00	1.50	
5	1.30	0.70		4.00	12.00		
6	1.24	0.72	2.47	6.00	4.00	1.50	
7	0.90	0.62	1.84	4.00	24.00	1.50	
8	1.04	0.82	2.14	10.00	12.00.	0.50	
9	0.97	0.76	2.00	4.00	14.00	1.50	
10	1.05	0.75	2.62	4.00	6.00	1.00	
11	0.90	0.75	1.75	14.00	14.00	4.00	
12	1.05	1.07	2.48	6.00	12.00	1.50	
13	1.62	0.90	1.85	4.03	24.00	4.00	
14	1.18	0.83	2.12	6.00	8.00	2.00	
15	0.88	0.63	2.11	4.00	12.02	1.00	
16	1.25	1.08	2.71	4.00	4.00	1.50	
17	1.01	0.80	2.50	4.00	6.00	0.50	
18	1.42	0.97	2.11	6.00	14.00	2.00	
N	18	18	17	18	18	17	
Mean	1.15	0.85	2.32	6.11	12.11	1.68	
SD	0.20	0.17	0.38	3.18	6.66	1.01	
Min	0.88	0.62	1.75	4.00	4.00	0.50	
Median	1.15	0.82	2.24	4.02	12.00	1.50	
Max	1.62	1.27	3.16	14.00	24.00	4.00	

- 5 C_{max} is the maximum measured plasma concentration of N-(2-propylpentanoyl) glycine after administration. T_{max} is the time at which the maximum concentration of N-(2-propylpentanoyl) glycine was measured.
- 10 N-(2-propylpentanoyl) glycine is one of the major metabolites of N- (2-propylpentanoyl) glycamine.

As seen in figure 2, formulations A and B maintain a mean plasma concentration of N-(2-propylpentanoyl) glycine which is stable from 4 hours after administration to 16 hours after administration. In addition, in formulations A and B, mean T_{max} occurs after 6 hours, whereas in formulation C, mean T_{max} occurs before 2 hours.

As seen in table 28, the C_{max} after administration of formulations A and B did not exceed 1.62 µg/ml in any of the volunteers. However, the mean C_{max} after administration of formulation C was 3.16 µg/ml. Administration of formulations A or B may eliminate unwanted side-effects which are caused as a result of dosage peaks present in immediate release formulations such as formulation C.

15

Discussion

In humans, neuropathic pain tends to be chronic. The same is true for epilepsy. In addition, epilepsy and neuropathic pain are diseases that require long term therapy. For most of the established drugs currently available for the treatment of these diseases, the required dosage must be administered several times daily. This results in compliance problems and fluctuations in plasma concentrations, which may lead to subtherapeutic and potentially toxic levels of the drug.

25 Development of sustained release formulations of anti neuropathic pain drugs and antiepileptic agents may improve the therapy of epileptic and/or neuropathic pain patients. The sustained release formulations of the present invention satisfy this pressing need.

30

In the present invention, the hydroxypropyl methyl cellulose is not part of the granule composition but is compressed with the granules into the final controlled release tablet. The formulations of the subject invention have the distinct advantage of allowing one to vary the desired dissolution

profile of the resulting tablet without requiring one to remake the granule composition. Thus, according to the present invention, one can manufacture granules of the active material in bulk and then vary the dissolution profile of the resulting tablets by varying the amount and type of hydroxypropyl methyl cellulose added to the mixture. In addition, the present invention does not require that specific sizes of the granules be selected for the resulting tablets. Consequently, the process of manufacture presented above is significantly easier to implement than a process in which the hydroxypropylmethyl cellulose is part of the granule composition.

In addition, as described earlier, N-(2-Propylpentanoyl)glycinamide is difficult to work with due to its "lamination and compression" characteristics. Thus, the subject invention provides the unexpected result of using a filler and two types of hydroxypropylmethyl cellulose to improve the compression characteristics while simultaneously slowing down the drug release profile. As illustrated in Example 4, the tablets manufactured according to the subject invention are also extremely stable.

Furthermore, as illustrated in the examples, the use of two types of hydroxypropylmethyl cellulose yields tablets which 25 release the drug at a steady rate over time, yet another advantage of the formulations of the subject invention.

Although the plasma concentration results in Example 11 are all based on administration of a single, 1000 mg dose of N-(230 propylpentanoyl) glycinamide, a linear pharmacokinetic response is expected in patients upon administration of other doses of similar formulation. Such a response is expected based on the work of Blotnick et al. with related compounds in phase I studies in which the pharmacokinetics were shown to be dose35 independent (Blotnick et al., "The Disposition of Valproyl

Glycinamide and Valproyl Glycine in Rats" (1997) Pharmaceutical Research 14(7): 873-878).

What is claimed is:

1. A sustained release solid dosage form comprising the following components:

- a) a uniform admixture of:
 - (i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$N$$
 $(CH_2)n$
 NR_2R_3

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

- (ii) a binder; and
- b) a hydroxypropylmethyl cellulose.

The solid dosage form of claim 1, wherein the solid dosage form is a tablet.

- 3. The solid dosage form of claim 1 or 2, wherein the uniform admixture of component a) further comprises a filler.
- 4. The solid dosage form of claim 3, wherein the filler comprises a microcrystalline cellulose.
- 5. The solid dosage form of claim 1 or 2, wherein the hydroxypropylmethyl cellulose comprises 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxyproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve, has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C, and has a pH in the range 5.5-8.0.
- 6. The solid dosage form of claim 5, wherein at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.
- 7. The solid dosage form of claim 1 or 2, further comprising as additional components a filler, a lubricant and a flow agent.
- 8. The solid dosage form of claim 1 or 2, wherein the binder of component a)(ii) comprises hydroxypropyl cellulose.

9. The solid dosage form of claim 1 or 2, further comprising a different hydroxypropylmethyl cellulose as a component.

- 10. The solid dosage form of claim 3, further comprising as additional components a filler, a lubricant and a flow agent.
- 11. The solid dosage form of claim 10, further comprising a different hydroxypropylmethyl cellulose as a component.
- 12. The solid dosage form of claim 9 or 11, wherein the different hydroxypropylmethyl cellulose comprises 19-24% by weight methoxyl substituent, 7-9% by weight hydroxypropoxyl substituent, has an apparent of 6,138-9,030 millipascal-seconds viscosity (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C, has a pH in the range 5.5-8.0 and has a particle size least distribution such that at 99% of hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.
- 13. The solid dosage form of claim 12, wherein at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.
- 14. The solid dosage form of claim 7, wherein
 the filler comprises a microcrystalline
 cellulose, anhydrous dicalcium phosphate, lactose,
 methylcellulose, carboxymethylcellulose, calcium
 carbonate, calcium sulfate kaolin, sodium chloride,

powdered cellulose, sucrose, mannitol or a combination of two or more of the foregoing;

the lubricant comprises magnesium stearate, sodium stearyl fumarate, hydrogenated castor oil, hydrogenated soybean oil, polyethylene glycol or a combination of two or more of the foregoing; and

the flow agent comprises a colloidal fumed silica, or colloidal silicon dioxide.

15. The solid dosage form of claim 14 wherein

the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant comprises magnesium stearate, sodium stearyl fumarate or a combination thereof; and

the flow agent comprises a colloidal fumed silica.

16. The solid dosage form of claim 1 or 2 wherein the active ingredient is a compound having the structure:

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3.

- 17. The solid dosage form of claim 16, wherein the active ingredient is N-(2-Propylpentanoyl)glycinamide.
- 18. A sustained release solid dosage form comprising the following components:
 - a) a uniform admixture of:
 - (i) N-(2-Propylpentanoyl)glycinamide; and
 - (ii) a binder;
 - b) a hydroxypropylmethyl cellulose; and
 - c) a different hydroxypropylmethyl cellulose.
- 19. The solid dosage form of claim 18, wherein the solid dosage form is a tablet.
- 20. The solid dosage form of claim 18 or 19, comprising a filler, a lubricant and a flow agent as additional components and wherein the uniform admixture of component a) further comprises a filler.
- 21. The solid dosage form of claim 20, wherein

the binder of component a)(ii) comprises hydroxypropyl cellulose;

the filler of component a) comprises a microcrystalline cellulose;

the hydroxypropylmethyl cellulose of component b) has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 'cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;

the hydroxypropylmethyl cellulose of component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;

the filler component comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant component comprises magnesium stearate, sodium stearyl fumarate or a combination thereof; and

the flow agent component comprises a colloidal fumed silica.

- 22. The solid dosage form of claim 21, comprising the following components:
 - a) a uniform admixture of:
 - (i) from 50 mg/solid dosage form to 1000 mg/solid dosage form of N-(2-propylpentanoyl) glycinamide,
 - (ii) from 1 mg/solid dosage form to 100 mg/solid dosage form hydroxypropyl cellulose; and

(iii)from 1 mg/solid dosage form to 200
mg/solid dosage form microcrystalline cellulose;

- b) from 10 mg/solid dosage form to 300 mg/solid dosage form of hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
- c) from 10 mg/solid dosage form to 300 mg/solid dosage form of a different hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
- d) from 1 mg/solid dosage form to 300 mg/solid dosage form microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
- e) from 0.1 mg/solid dosage form to 20 mg/solid dosage form of magnesium stearate, sodium stearyl fumarate or a combination thereof; and f) from 0.1 mg/solid dosage form to 15 mg/solid dosage form a colloidal fumed silica.
- 23. The solid dosage form of claim 21, comprising the following components:
 - a) a uniform admixture of:
 - (i) from 500 mg/solid dosage form to 850 mg/solid dosage form of N-(2-propylpentanoyl) glycinamide,

(ii) from 25 mg/solid dosage form to 75 mg/solid dosage form hydroxypropyl cellulose; and

- (iii) from 50 mg/solid dosage form to 150 mg/solid dosage form microcrystalline cellulose;
- b) from 100 mg/solid dosage form to 300 mg/solid dosage form of hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
- c) from 20 mg/solid dosage form to 150 mg/solid dosage form of a different hydroxypropylmethyl cellulose having 19%-24% by weight methoxyl substituent, 7%-12% hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;
- d) from 20 mg/solid dosage form to 100 mg/solid dosage form microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
- e) from 2 mg/solid dosage form to 20 mg/solid dosage form of magnesium stearate, sodium stearyl fumarate or a combination thereof; and
- f) from .5 mg/solid dosage form to 5 mg/solid dosage form a colloidal fumed silica, per 1 gram solid dosage form.
- 24. The solid dosage form of any one of claims 22 or 23, wherein at least 90% of the hydroxypropylmethyl cellulose of component b), of component c), or of

both component b) and c) passes through a No. 100 US standard sieve.

25. The solid dosage form of claim 23, wherein

the hydroxypropylmethy1 cellulose of component b) has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the hydroxypropylmethyl cellulose of component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

- 26. The solid dosage form of claim 23, comprising the following components:
 - a) a uniform admixture of :
 - (i) 500 mg/solid dosage form N-(2-Propylpentanoyl)glycinamide,
 - (ii) 50 mg/solid dosage form hydroxypropyl
 cellulose; and
 - (iii) 100 mg/solid dosage form microcrystalline cellulose;
 - b) 150 mg/solid dosage form of hydroxypropylmethylcellulose having 19%-24% by weight methoxyl
 substituent, 7%-12% by weight hydroxylproproxyl
 substituent and has a particle size distribution
 such that at least 99% of the hydroxypropylmethyl
 cellulose passes through a No. 40 US standard
 sieve;
 - c) 60 mg/solid dosage form of a different hydroxypropylmethyl cellulose having 19%-24% by

weight methoxyl substituent, 7%-12% hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve;

- d) 20 mg/solid dosage form lactose;
- e) 4.5 mg/solid dosage form magnesium stearate; and
- f) 1 mg/solid dosage form colloidal fumed silica.
- 27. The solid dosage form of claim 26, wherein at least 90% of the hydroxypropylmethyl cellulose of component b), of component c), or of both component b) and c) passes through a No. 100 US standard sieve.
- 28. The solid dosage form of claim 26, wherein

the hydroxypropylmethyl cellulose of component b) has an apparent viscosity of 78-117 millipascalseconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the hydroxypropylmethyl cellulose of component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

- 29. A hard compressed tablet comprising a uniform admixture of the following components:
 - a) N-(2-Propylpentanoyl)glycinamide;
 - b) a hydroxypropylmethyl cellulose; and
 - c) a different hydroxypropylmethyl cellulose.

30. The tablet of claim 29, wherein

the hydroxypropylmethyl cellulose component b) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve; and

the hydroxypropylmethyl cellulose component c) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

- 31. The tablet of any one of claims 29 or 30, wherein at least 90% of the hydroxypropylmethyl cellulose of component b), of component c), or of both component b) and c) passes through a No. 100 US standard sieve.
- 32. The tablet of claim 30, wherein

the hydroxypropylmethyl cellulose component b) has an apparent viscosity of 78-117 millipascalseconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the hydroxypropylmethyl cellulose component c) has an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

33. The tablet of claim 29, further comprising a filler, lubricant and flow agent as additional components.

- 34. The tablet of claim 33, wherein
 - the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant comprises sodium stearyl fumarate; and

the flow agent comprises a colloidal fumed silica.

- 35. The tablet of claim 34, comprising a uniform admixture of the following components:
 - a) from 100 mg/tablet to 1000 mg/tablet N-(2-Propylpentanoy1)glycinamide;
 - b) from 10 mg/tablet to 300 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;
 - c) from 10 mg/tablet to 300 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;
 - d) from 1 mg/tablet to 300 mg/tablet a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

e) from 0.1 mg/tablet to 20 mg/tablet sodium stearyl fumarate; and

- f) from 0.1 mg/tablet to 15 mg/tablet a colloidal fumed silica.
- 36. The tablet of claim 34, comprising a uniform admixture of the following components:
 - a) from 400 mg/tablet to 1000 mg/tablet N-(2-Propylpentanoyl)glycinamide;
 - b) from 100 mg/tablet to 300 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;
 - c) from 20 mg/tablet to 150 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;
 - d) from 10 mg/tablet to 60 mg/tablet a microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;
 - e) from 2 mg/tablet to 20 mg/tablet sodium stearyl fumarate; and
 - f) from 5 mg/tablet to 15 mg/tablet a colloidal fumed silica, per 1 gram tablet.
- 37. The tablet of claim 36, comprising a uniform admixture of the following components:

a) 500 mg/tablet N-(2-Propylpentanoyl)glycinamide;

- b) 150 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascalseconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C;
- c) 60 mg/tablet of hydroxypropylmethyl cellulose with an apparent viscosity of 6,138-9,030 millipascalseconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C;
- d) 20 mg/tablet lactose;
- e) 10 mg/tablet sodium stearyl fumarate; and
- f) 10 mg/tablet colloidal fumed silica.
- 38. A composition in granulate form comprising a uniform admixture of:
 - (i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

$$\bigcap_{\mathbf{H}} \bigcap_{(CH_2)_{\Pi}} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

- (ii) a hydroxypropyl cellulose.
- 39. The composition of claim 38, wherein the active ingredient comprises a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N \text{ (CH}_2)m} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a $C_1\text{--}C_6$ alky1 group, an

aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3.

- 40. The composition of claim 38, wherein the active ingredient comprises valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium or valpromide.
 - 41. A tablet comprising the granulate of claim 38 as a component.
 - 42. The tablet of claim 41, wherein the granulate further comprises a filler.
 - 43. The tablet of claim 41, further comprising a hydroxypropylmethyl cellulose as a component.
 - 44. The tablet of claim 41, further comprising as additional components a filler, a lubricant and a flow agent.
 - 45. The tablet of claim 43, further comprising as additional components a filler, a lubricant and a flow agent.
 - 46. The tablet of claim 43, further comprising a different hydroxypropylmethyl cellulose as a component.
 - 47. The tablet of claim 43, wherein

the hydroxypropylmethyl cellulose has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle

size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

48. The tablet of claim 47, wherein at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

49. The tablet of claim 47, wherein

the hydroxypropylmethyl cellulose has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C.

50. The tablet of claim 46, wherein

the different hydroxypropylmethyl cellulose. has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

51. The tablet of claim 50, wherein at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

52. The tablet of claim 50, wherein

the different hydroxypropylmethyl cellulose has an apparent viscosity of 6,138-9,030 millipascalseconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

53. The tablet of claim 42, wherein the filler in the granulate is a microcrystalline cellulose.

54. The tablet of claim 45, wherein

the filler comprises a microcrystalline cellulose, anhydrous dicalcium phosphate,

lactose or a combination of two or more of the foregoing;

the lubricant comprises magnesium stearate, sodium stearyl fumarate or a combination thereof; and

the flow agent comprises a colloidal fumed silica.

55. A sustained release tablet comprising a compound having the structure:

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer

which is greater than or equal to 0 and less than or equal to 3.

- 56. The sustained release tablet of claim 55, wherein the compound is N-(2-propylpentanoyl)glycinamide.
- 57. A method of treating neuropathic pain in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby treat the neuropathic pain in the subject.
- 58. A method of treating a headache disorder in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby treat the headache disorder in the subject.
- 59. A method of treating epilepsy in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby treat epilepsy in the subject.
- 60. A method of controlling seizures in a subject suffering from epilepsy comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the

tablet of any one of claims 29-37 or 41-56 in order to thereby control the seizures in the subject.

- 61. A method of treating pain in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby treat pain in the subject.
- 62. A method of pain prophylaxis in a subject in need of such treatment comprising administering to the subject a prophylactic dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby effect pain prophylaxis in the subject.
- 63. A method of treating mania in bipolar disorder in a subject in need of such treatment comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby treat mania in bipolar disorder in the subject.
- 64. A method of attenuating bipolar mood swings in a subject suffering from bipolar disorder comprising administering to the subject a therapeutically effective dose of the solid dosage form of any one of claims 1-28 or the tablet of any one of claims 29-37 or 41-56 in order to thereby attenuate the bipolar mood swings in the subject.

65. A process for preparing the solid dosage form of claim 1 or 2, comprising the steps of:

- a) admixing predetermined amounts of
 - (i) an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$N$$
 (CH_2)
 N
 NR_2R_3

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3; and

(ii) a binder;

- b) admixing the uniform mixture of step a) with a predetermined amount of a hydroxypropylmethyl cellulose; and
- c) compressing the mixture of step b) to form the tablet.

66. The process of claim 65, wherein step b) further comprises admixing the uniform mixture with a predetermined amount of a different hydroxypropylmethyl cellulose.

- 67. The process of claim 66, wherein step b) further comprises admixing the uniform mixture with predetermined amounts of a filler, a lubricant and a flow agent.
- 68. The process of claim 67, wherein the flow agent comprises colloidal fumed silica.
- 69. The process of claim 67, wherein the filler comprises microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing.
- 70. The process of claim 69, wherein the filler comprises lactose.
- 71. The process of claim 67, wherein the lubricant comprises magnesium stearate or sodium stearyl fumarate or a combination thereof.
- 72. The process of claim 71, wherein the lubricant comprises magnesium stearate.
- 73. The process of claim 66, wherein

each hydroxypropylmethyl cellulose of step b) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of

the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

74. The process of claim 73, wherein at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

75. The process of claim 73, wherein

the first hydroxypropylmethyl cellulose has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the second hydroxypropylmethyl cellulose has an apparent viscosity of 6,138-9,030 millipascalseconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

- 76. A process for preparing the hard compressed tablet of claim 29 comprising the steps of:
 - a) admixing predetermined amounts of N-(2-Propylpentanoy1)glycinamide, hydroxypropylmethy1 cellulose, and a different hydroxypropylmethyl cellulose; and
 - b) compressing the mixture of step a) to form the hard compressed tablet.
- 77. The process of claim 76, wherein

each hydroxypropylmethyl cellulose of step a) has 19%-24% by weight methoxyl substituent, 7%-12% by weight hydroxylproproxyl substituent and has a particle size distribution such that at least 99% of

the hydroxypropylmethyl cellulose passes through a No. 40 US standard sieve.

78. The process of claim 77, wherein at least 90% of the hydroxypropylmethyl cellulose passes through a No. 100 US standard sieve.

79. The process of claim 77, wherein

the hydroxypropylmethyl cellulose has an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and

the different hydroxypropylmethyl cellulose has an apparent viscosity of 6,138-9,030 millipascalseconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C.

- 80. The process of claim 76, wherein step a) further comprises admixing predetermined amounts of a filler, lubricant and flow agent as additional components.
- 81. The process of claim 80, wherein

the filler comprises microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing;

the lubricant comprises sodium stearyl fumarate; and

the flow agent comprises colloidal fumed silica.

82. A process for preparing the composition in granulate form of claim 38, comprising granulating a predetermined amount of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide or a compound having the structure:

$$\bigcap_{N \in \mathbb{C}(CH_2)} \bigcap_{N \in \mathbb{R}_2 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3 \setminus \mathbb{R}_3} \bigcap_{N \in \mathbb{R}_3} \bigcap_{$$

or

$$\bigcap_{N} \bigcap_{(CH_2)\pi} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, and a predetermined amount of hydroxypropyl cellulose to form the composition in granulate form.

- 83. A process for preparing a sustained release tablet comprising the steps of:
 - a) admixing the granules of claim 38 with predetermined amounts of a hydroxypropylmethyl cellulose; and

b) compressing the mixture of step a) to form the tablet.

- 84. The process of claim 83, wherein step a) further comprises admixing the granules with a predetermined amount of each of a different hydroxypropylmethyl cellulose, a filler, a lubricant and a flow agent.
- 85. The process of claim 84, wherein the flow agent comprises colloidal fumed silica.
- 86. The process of claim 84, wherein the filler comprises microcrystalline cellulose, anhydrous dicalcium phosphate, lactose or a combination of two or more of the foregoing.
- 87. The process of claim 86, wherein the filler is lactose.
- 88. The process of claim 84, wherein the lubricant comprises magnesium stearate or sodium stearyl fumarate or a combination thereof.
- 89. The process of claim 88, wherein the lubricant comprises magnesium stearate.
- 90. The process of claim 83, comprising the steps of:

 a) admixing the granules with predetermined amounts of hydroxypropyl methyl cellulose having an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°C, and

hydroxypropyl methyl cellulose having an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C; and b) compressing the mixture of step a) to form the tablet.

- 91. The process of claim 90, wherein step a) further comprises admixing the granules with predetermined amounts of a flow agent, a filler, and a lubricant.
- 92. The process of claim 91 comprising the steps of a) admixing the granules with
 - a predetermined amount of hydroxypropylmethyl cellulose with an apparent viscosity of 78-117 millipascal-seconds (nominal value 98 mPa.s) by rotation and 80-120 cP (nominal value 100 cP) by Ubbelhode, at a concentration of 1% by weight in water at 20°C which results in tablets containing 150 mg/tablet;
 - a predetermined amount of hydroxypropyl methyl cellulose with an apparent viscosity of 6,138-9,030 millipascal-seconds (nominal value 7382 mPa.s) by rotation and 11,250-21,000 cP (nominal value 15,000 cP) by Ubbelhode at a concentration of 1% by weight in water at 20°Cwhich results in tablets containing 60 mg/tablet;
 - a predetermined amount of lactose which results in tablets containing 20 mg/tablet;
 - a predetermined amount of magnesium stearate which results in tablets containing 4.5 mg/tablet; and

a predetermined amount of a colloidal fumed silica which results in tablets containing 1 mg/tablet; and

- b) compressing the mixture of step a) to form the tablet.
- O3. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH2)n}} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating a headache disorder in a subject.

94. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

or

$$N$$
 $(CH_2)n$
 NR_2R_3

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating neuropathic pain in a subject.

95. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N \text{ } (CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating epilepsy in a subject.

96. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in controlling seizures in a subject suffering from epilepsy.

97. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating mania in bipolar disorder in a subject.

98. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N \in \mathbb{N}} \mathbb{N} = \{ (CH_2)n \cap \mathbb{N} \}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in attenuating bipolar mood swings in a subject suffering from bipolar mood disorder.

99. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N} \bigcap_{\text{CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$\bigcap_{N} \bigcap_{(CH_2)n} \bigcap_{NR_2R_3}$$

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or

equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating pain in a subject.

100. Use of an active ingredient selected from the group consisting of valproic sodium acid, a pharmaceutically acceptable salt or ester of valproic acid, divalproex sodium, valpromide and a compound having the structure:

$$\bigcap_{N \text{ (CH}_2)n} \bigcap_{NR_2R_3}$$

or

$$N$$
 $(CH_2)n$
 NR_2R_3

wherein R_1 , R_2 , and R_3 are independently the same or different and are hydrogen, a C_1 - C_6 alkyl group, an aralkyl group, or an aryl group, and n is an integer which is greater than or equal to 0 and less than or equal to 3, for manufacturing a sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in effecting pain prophylaxis in a subject.

PCT/US2004/003281 WO 2004/071421

101. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating a headache disorder in a subject.

~ ** ·

- 102. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating neuropathic pain in a subject.
- 103. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating epilepsy in a subject.
- 104. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in controlling seizures in a subject suffering from epilepsy.
- 105. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating mania in bipolar disorder in a subject.
- 105. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in attenuating bipolar mood swings in a subject suffering from bipolar disorder.
- of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in treating pain in a subject.

107. The sustained release solid dosage form of any one of claims 1-28 or tablet of any one of claims 29-37 or 41-56 for use in effecting pain prophylaxis in a subject.

- 108. A controlled release oral unit dose composition comprising N-(2-propylpentanoyl) glycinamide and at least one pharmaceutically acceptable carrier, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 4 and 24 hours after ingestion of a single oral unit dose.
- 109. The controlled release oral unit dose composition of claim 108, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 4 and 12 hours after ingestion of a single oral unit dose.
- 110. The controlled release oral unit dose composition of claim 109, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 6 and 12 hours after ingestion of a single oral unit dose.
- 111. The controlled release oral unit dose composition of claim 110, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoy1) glycinamide between 6 and 8 hours after ingestion of a single oral unit dose.

PCT/US2004/003281 WO 2004/071421

112. The controlled release oral dose composition of any one of claims 108 to 111, wherein the peak blood plasma level of N-(2-propylpentanoyl) glycinamide is from 0.5 micrograms/ml to 16 micrograms/ml per a 1000 mg dose of N-(2-propylpentanoyl) glycinamide in the composition.

- 113. The controlled release oral dose composition of claim 108, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2-propylpentanoyl) glycine in the human subject from 0.5 $\mu g/mL$ to 1.7 $\mu g/mL$ per a 1000 mg dose of N-(2-propylpentanoyl) glycinamide in the composition.
 - composition dose oral release controlled comprising N-(2-propylpentanoy1) glycinamide and a 114. A wherein the pharmaceutically acceptable carrier, composition when orally ingested by a human subject, induces a peak blood plasma level propylpentanoyl) glycinamide of 0.5 μ g/mL to 16 $\mu g/mL$ per a 1000 mg dose in the composition.
 - composition dose oral release controlled comprising N-(2-propylpentanoyl) glycinamide and a 115. A pharmaceutically acceptable carrier, wherein the composition when orally ingested by a human subject, induces a peak blood plasma level of N-(2propylpentanoyl) glycine of 0.5 $\mu g/mL$ to 1.7 $\mu g/mL$ dose of N-(2-propylpentanoy1) 1000 mg glycinamide in the composition.
 - 116. A method of inducing in a human subject a peak blood plasma level of N-(2-propylpentanoy1) glycinamide

between 4 and 24 hours after administration of N-(2-propylpentanoyl) glycinamide, comprising administering to the human subject a controlled release oral unit dose composition comprising N-(2-propylpentanoyl) glycinamide and at least one pharmaceutically acceptable carrier, which composition induces a peak blood plasma level of N-(2-propylpentanoyl) glycinamide between 4 and 24 hours after administration of a single oral unit dose.

- 117. The method of claim 116, wherein the peak blood plasma level of N-(2-propylpentanoyl) glycinamide occurs between 4 and 12 hours after administration.
- 118. The method of claim 116, wherein the peak blood plasma level of N-(2-propylpentanoyl) glycinamide is 0.5 μ g/mL to 16 μ g/mL per 1000 mg dose of N-(2-propylpentanoyl) glycinamide in the composition.
- 119. The method of any one of claims 116-118, wherein the administration to the human subject of a controlled release oral unit dose composition comprising N-(2-propylpentanoyl) glycinamide and at least one pharmaceutically acceptable carrier induces a peak blood plasma level of N-(2-propylpentanoyl) glycine in the human subject from 0.5 µg/mL to 1.7 µg/mL upon administration of a single 1000 mg dose of N-(2-propylpentanoyl) glycinamide.
- 120. The method of any one of claims 116-119, wherein the controlled release oral dose composition is the solid dosage form of any one of claims 18-28 or the tablet of any one of claims 29-37 or 41-56.

ägure l

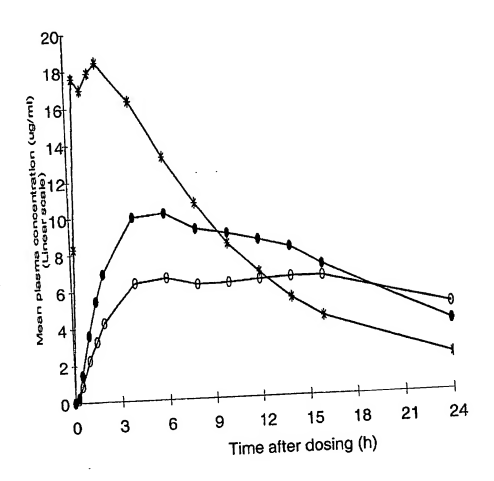
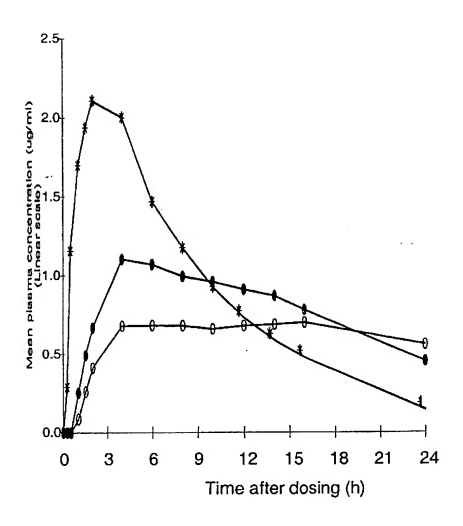


Figure 2



THIS PAGE BLANK (USPTO)